



THE UNIVERSITY *of* EDINBURGH

## Edinburgh Research Explorer

### Enhancing nano-scale computational fluid dynamics with molecular pre-simulations: unsteady problems and design optimisation

**Citation for published version:**

Holland, DM, Borg, MK, Lockerby, DA & Reese, JM 2015, 'Enhancing nano-scale computational fluid dynamics with molecular pre-simulations: unsteady problems and design optimisation', *Computers and Fluids*, vol. 115, pp. 46-53. <https://doi.org/10.1016/j.compfluid.2015.03.023>

**Digital Object Identifier (DOI):**

[10.1016/j.compfluid.2015.03.023](https://doi.org/10.1016/j.compfluid.2015.03.023)

**Link:**

[Link to publication record in Edinburgh Research Explorer](#)

**Document Version:**

Publisher's PDF, also known as Version of record

**Published In:**

Computers and Fluids

**General rights**

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

**Take down policy**

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact [openaccess@ed.ac.uk](mailto:openaccess@ed.ac.uk) providing details, and we will remove access to the work immediately and investigate your claim.





# Enhancing nano-scale computational fluid dynamics with molecular pre-simulations: Unsteady problems and design optimisation



David M. Holland<sup>a</sup>, Matthew K. Borg<sup>b</sup>, Duncan A. Lockerby<sup>a,\*</sup>, Jason M. Reese<sup>c</sup>

<sup>a</sup> School of Engineering, University of Warwick, Coventry CV4 7AL, United Kingdom

<sup>b</sup> Department of Mechanical & Aerospace Engineering, University of Strathclyde, Glasgow G1 1XJ, United Kingdom

<sup>c</sup> School of Engineering, University of Edinburgh, Edinburgh EH9 3JL, United Kingdom

## ARTICLE INFO

### Article history:

Received 31 December 2014

Received in revised form 27 February 2015

Accepted 26 March 2015

Available online 1 April 2015

### Keywords:

Nanofluidics

Computational fluid dynamics

Molecular dynamics

Hybrid methods

Design optimisation

Murray's Law

## ABSTRACT

We demonstrate that a computational fluid dynamics (CFD) model enhanced with molecular-level information can accurately predict unsteady nano-scale flows in non-trivial geometries, while being efficient enough to be used for design optimisation. We first consider a converging–diverging nano-scale channel driven by a time-varying body force. The time-dependent mass flow rate predicted by our enhanced CFD agrees well with a full molecular dynamics (MD) simulation of the same configuration, and is achieved at a fraction of the computational cost. Conventional CFD predictions of the same case are wholly inadequate. We then demonstrate the application of enhanced CFD as a design optimisation tool on a bifurcating two-dimensional channel, with the target of maximising mass flow rate for a fixed total volume and applied pressure. At macro scales the optimised geometry agrees well with Murray's Law for optimal branching of vascular networks; however, at nanoscales, the optimum result deviates from Murray's Law, and a corrected equation is presented.

© 2015 The Authors. Published by Elsevier Ltd. This is an open access article under the CC BY license (<http://creativecommons.org/licenses/by/4.0/>).

## 1. Introduction

Many emerging applications of nanofluidic technology take advantage of different physical effects that dominate at small scales; examples can be found in air and water purification [1,2], and in micro chemical reactors [3,4]. The design of these technologies would be greatly facilitated by being able to perform numerical simulations that predict mass flow rates and heat transfer. Computational fluid dynamics (CFD) is regularly used to model and create optimal every-day engineering designs efficiently. However, the assumptions used to derive the continuum fluid equations become invalid in highly-confined systems, making the equations inaccurate. On the other hand, molecular dynamics (MD) can be used to perform highly detailed simulations of nano-scale systems; it has been successfully used to study the behaviour of protein folding [5], crystal formation [6] and chemical reactions [7]. The drawback is that MD is extremely computationally intensive, especially when used to model systems comprising hundreds of thousands of molecules that would be required for engineering applications.

The continuum fluid assumptions become inaccurate for gas flows as the smallest characteristic scale of the geometry (e.g.

channel height) approaches the mean distance between molecular collisions (i.e. the mean free path) [8]. When modelling dense liquids (as we do in this paper) there is not a well-defined condition for when the fluid assumptions become inaccurate. However, it appears that they fail when water is confined in channels of width  $\sim 1\text{--}2\text{ nm}$  (see [9] and references therein), and MD simulations have been used to show that Lennard-Jones fluids confined in geometries of  $\sim 2\text{--}3\text{ nm}$  still show continuum behaviour [10–12]. At the nano-scale the fluid molecules form layers parallel to an interface, which causes the strain rate to vary rapidly within several molecular diameters [13]. These large variations mean the stress no longer has local linear behaviour [14,15].

Despite the complexities of fluid behaviour at the nano-scale, it has recently been shown that useful predictions from CFD can be obtained for some simple geometries, if appropriate fluid state models, viscosity relationships, and slip models are extracted from an MD pre-simulation [16]. In [17], a similar approach is used to obtain CFD predictions of flow through a nanotube; with results agreeing well with full MD simulations.

However, what remains to be tested is the robustness of nano-scale CFD when applied to more complex engineering calculations. In this paper, we test if our enhanced CFD model is robust enough to predict flow behaviour in a non-trivial geometry (a converging–diverging channel), while using various forms of applied forcing to generate unsteadiness within it. As a demonstration of its

\* Corresponding author.

E-mail address: [duncan.lockerby@warwick.ac.uk](mailto:duncan.lockerby@warwick.ac.uk) (D.A. Lockerby).

efficiency, we go on to apply the enhanced CFD to the design optimisation of a small fluidic network.

The paper is structured as follows. In the next section we summarise the MD pre-simulation procedure and the CFD model used. We then use these models to perform unsteady simulations of a converging–diverging channel, where the width of the channel is close to the continuum limit. Our model is then applied to simulate an industrially-relevant problem of flow through a bifurcating channel. We show that to optimally design the channels the slip velocity at walls must be taken into account. The paper then ends with a summary.

## 2. Methodology

### 2.1. The MD pre-simulations

As in [16], we employ preliminary molecular dynamics (MD) simulations to obtain fluid properties and boundary conditions that enable the effective use of a Navier–Stokes fluid solver for nano-scale applications. This approach can be classed as a “sequential molecular-continuum hybrid method” (see [18] for a review of hybrid methods), where ‘sequential’ refers to the fact that the MD is performed in advance of, and so independent of, the continuum-fluid solver.

Fig. 1 (far left) shows a schematic of the MD pre-simulation domain; it is symmetrical about its centrelines and uses periodic boundary conditions in the streamwise direction (in the  $x$ -direction) and into the page (in the  $z$ -direction). The domain has *bulk*, *shear* and *interface* zones (as labelled) for measuring state, constitutive and boundary properties, respectively. Pressure and density are measured in the bulk zone. In addition to this, in the bulk zone an artificial streamwise body force ( $F_x$ ) is applied (Fig. 1, centre left), which creates a velocity profile in the domain similar to that illustrated (Fig. 1, centre right). We assume that the equation

of state in the bulk zone is unaffected by the magnitude of strain rate generated. In the shear zone the fluid is, therefore, subject to a constant shear stress,  $\tau_{xy}$ , directly resulting from the bulk-zone forcing. A linear flow velocity profile is developed in the shear zone, and this is least-squares fitted to obtain a strain rate and shear viscosity coefficient,  $\mu$ .

Any significant density oscillations associated with molecular layering are confined to the interface zone. In this zone we calculate what we term the ‘CFD surface displacement’,  $\delta$ , which is the distance that a CFD wall/surface needs to be displaced from the centres of surface atoms in order to accurately represent the boundary of the fluid (as opposed to the boundary of the solid); see  $\delta$  in Fig. 1. We take this displacement to be the distance from the centre of the surface wall atoms to where the fluid density becomes at least 10% of the bulk, i.e.  $\rho \geq a\rho_{\text{bulk}}$ , where  $a = 0.1$ . Note, the surface displacement is quite insensitive to the percentage of the bulk density chosen as the threshold, since the density increases from zero to well above the bulk density over a very short distance. For example, had we chosen the threshold to be at 20% of the bulk density, the surface displacement would have only been 1–2% larger, for a typical case.

The linear velocity profile obtained in the shear zone is extrapolated into the interface zone to find the apparent slip length,  $\xi$ , as defined from the CFD surface (see Fig. 1, centre right).

The molecular dynamics pre-simulations, and the full-scale MD simulations used for benchmarking, are performed using the *mdFoam* solver [19–22] that is implemented within the OpenFOAM libraries [23]. For the test cases considered in this paper we have adopted a simple Lennard-Jones (LJ) fluid model (at 292.8 K), where the solid LJ wall atoms are fixed/frozen [24]. However, the methodology is general to any given molecular model. For full details of the molecular pre-simulation domain and the molecular dynamics parameters used, the reader is referred to [16].

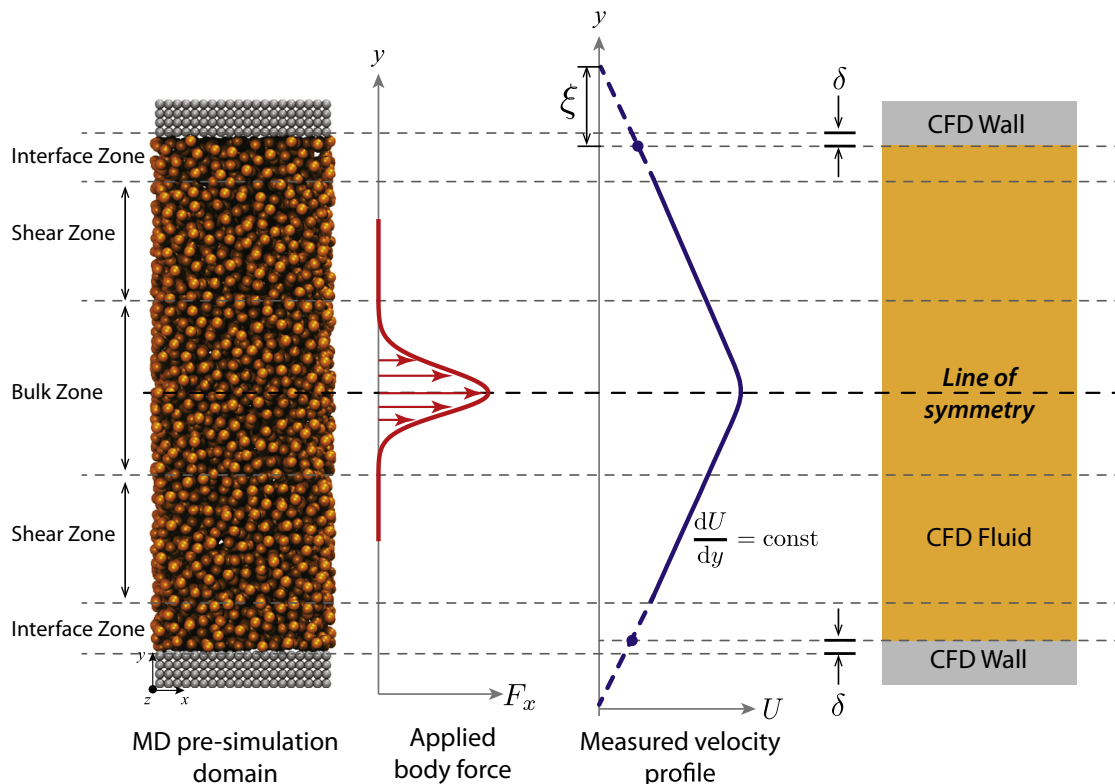


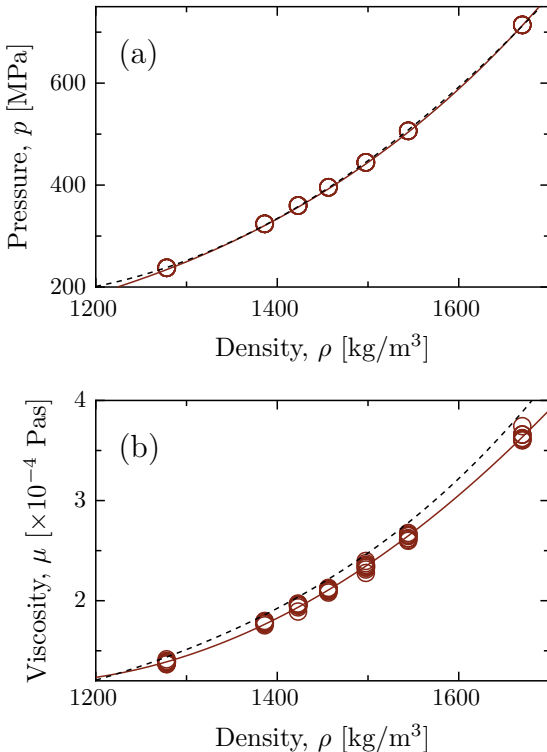
Fig. 1. Schematic of molecular dynamics pre-simulation for extracting fluid dynamic properties that are essential inputs to an enhanced CFD solver for nano-scale flows.

Fig. 2(a) shows MD pre-simulation measurements of pressure, obtained from the standard Irving–Kirkwood expression [25], varying with the mass density. The MD pre-simulation results are least-squares-fitted to a 2nd order polynomial. This then serves as an equation of state within the enhanced CFD solver to connect the mass continuity equation to the momentum equation. In this case the polynomial is  $p = 0.001559\rho^2 - 3.387\rho + 2020.6$ .

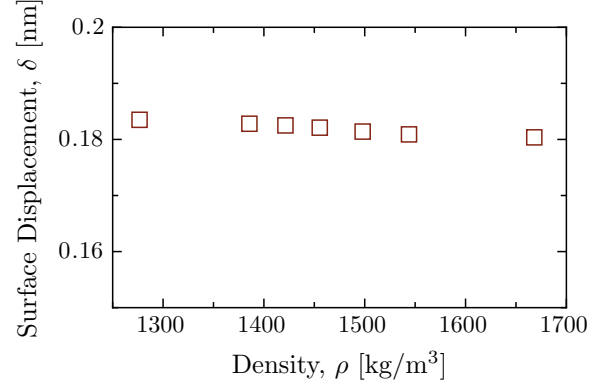
The strain-rate is extracted from the MD shear zone by a least-squares linear fit to the relaxed and time-averaged velocity profile. The applied shear stress is measured using the Irving–Kirkwood equation, from which we obtain a dynamic shear viscosity coefficient for the LJ fluid at a given bulk density. The viscosity coefficients measured from our MD pre-simulations of Lennard-Jones argon are shown in Fig. 2(b). A least-squares polynomial fit of 2nd order in density is also plotted:  $\mu = 7.96 \times 10^{-10}\rho^2 - 1.774 \times 10^{-6}\rho + 0.001106$ . This is then used in our enhanced CFD simulations to close the momentum equation. Note, due to the breakdown of the continuum assumption and the existence of non-local stress, this state-dependent viscosity becomes only approximate when applied to a nano-confined fluid.

The surface displacement  $\delta$  defines the location of the CFD boundaries relative to the atomic (actual) walls. If  $\delta$  varies substantially with density (or any other fluid property), the geometry of the enhanced CFD domain becomes dependent on the CFD solution itself. However, for the fluid/solid combination considered in this paper, over the density ranges considered,  $\delta$  is effectively constant, as seen in Fig. 3.

In certain cases the value of  $\delta$  will itself be dependent on the geometry, particularly for high curvatures, such as around sharp corners and obstructions. It is beyond the scope of the current work to attempt to accommodate these influences, while noting that, later, we obtain good agreement with full MD simulations



**Fig. 2.** Data for the Lennard-Jones fluid properties: (a) pressure variation with density, and (b) viscosity variation with density. MD data points from pre-simulation (circles), fitted polynomial (solid lines) and NIST data [26] (dashed lines).



**Fig. 3.** Surface displacement  $\delta$  varying with fluid density  $\rho$ , measured from each MD pre-simulation.

without doing so. To tackle geometry-dependent flow properties (including surface displacement) would dramatically increase the parameter space that the pre-simulations would be required to supply information for; in fact, for such problems a ‘concurrent’ hybrid approach is likely to be more efficient.

As the spatial-scale of the geometry increases, the relative significance of the surface displacement reduces. We can develop a simple gauge of its impact by considering the percentage that it modifies the mass flow rate in a simple channel in two limiting cases: assuming no-slip at the walls (i.e.  $\xi = 0$ ); and for very high slip (i.e.  $\xi \gg h$ , where  $h$  is the channel width). In the no-slip case, for Poiseuille flow, the mass flow rate is proportional to the cube of the channel width; the percentage difference of using the surface displacement is then

$$\epsilon = \left( 1 - \frac{(h - 2\delta)^3}{h^3} \right) \times 100\%. \quad (1)$$

For the cases in Section 3, where the channel width varies,  $\epsilon$  is  $\sim 28$ – $44\%$ . For high-slip cases, where the velocity profile becomes plug-like, the mass flow rate becomes proportional to the square of the channel width, giving a percentage difference:

$$\epsilon = \left( 1 - \frac{(h - 2\delta)^2}{h^2} \right) \times 100\%. \quad (2)$$

Considering again the cases in Section 3,  $\epsilon$  would be  $\sim 20$ – $32\%$ ; i.e. the impact of the surface displacement is likely to be very significant regardless of the degree of velocity slip. Based on the estimates of Eqs. (1) and (2), the impact of a surface displacement  $\delta \sim 0.2$  nm will only be less than 1% (i.e. negligible) for channels greater than 75–100 nm.

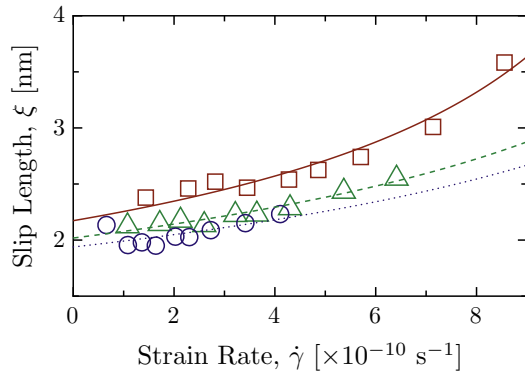
Liquid slip velocity at surfaces is calculated using the Navier slip condition:

$$u_{\text{slip}} = \xi \dot{\gamma}, \quad (3)$$

where  $\xi$  is the slip-length and  $\dot{\gamma}$  is the shear-rate at the bounding surface. The least-squares-fitted linear velocity profile is used to calculate the slip-length (as defined from the CFD surface). Based on the strain-rate/slip-length relationship proposed in [24], and assuming a linear dependence on density, a least-squares fit is performed to the following equation:

$$\xi = \frac{(c_1 \rho + c_2)}{\sqrt{1 - \dot{\gamma}/\dot{\gamma}_c}}, \quad (4)$$

where  $\rho$  is the density,  $\dot{\gamma}_c$  is the critical shear rate (see [24]), and  $c_1$ ,  $c_2$  and  $\dot{\gamma}$  are parameters of the fit to our MD pre-simulations, which are  $-1.205 \times 10^{-12} \text{ kg}^{-1} \text{ m}^4$ ,  $3.747 \times 10^{-9} \text{ m}$  and



**Fig. 4.** Slip length  $\xi$  varying with strain rate  $\dot{\gamma}$  for three density values  $\rho_1 = 1276 \text{ kg/m}^3$ ,  $\rho_2 = 1447 \text{ kg/m}^3$  and  $\rho_3 = 1668 \text{ kg/m}^3$ . MD pre-simulation data points (symbols) and fit to Eq. (4) (dashed lines).

$1.543 \times 10^{11} \text{ s}^{-1}$ , respectively. Fig. 4 shows our MD pre-simulation data and the least-squares fit of Eq. (4); data are shown for three different values of density. The slip model approximated by Eqs. (3) and (4) is directly introduced as a Robin boundary condition in the enhanced CFD solver.

## 2.2. The enhanced CFD model

We use the laminar, compressible flow solver *sonicLiquidFoam*, which we have modified to (a) accommodate a nonlinear equation of state, (b) allow a density-dependent viscosity, and (c) incorporate slip boundary conditions of the form given in Eq. (4). A compressible solver is used despite the very low Mach numbers, since significant compressibility can occur in micro and nano geometries due to very high viscous pressure losses [27,28].

## 3. Unsteady simulations

We now simulate the unsteady flow behaviour of a Lennard-Jones fluid along a converging-diverging channel; a case chosen to demonstrate the robustness of the enhanced CFD model when applied to non-trivial flow problems.

Owing to the lack of detailed and reliable experimental flow measurements at the nano-scale, in this section we compare our enhanced CFD predictions with full-scale MD simulation results. This comparison is intended to test whether enhanced CFD can produce flow field solutions of comparable accuracy to full MD in complex nano-scale geometries, without the need for *ad hoc* corrections, and at only a fraction of the cost of full MD.

### 3.1. Cases

We consider a two-dimensional geometry: a converging-diverging channel with a smoothly varying height in the streamwise direction. A gravity-type force is applied to the fluid to generate an unsteady/transient flow. As test cases, we choose flows that exhibit non-continuum behaviour (e.g. slip at surfaces), and do not contain a significant bulk flow region, i.e., the width of the channel is at the 2–3 nm continuum-fluid limit for a Lennard-Jones fluid.

The converging-diverging channel is shown in Fig. 5 and has a length  $l = 68 \text{ nm}$  in the streamwise direction  $x$ , a depth of  $5.44 \text{ nm}$ , and heights of  $3.4 \text{ nm}$  and  $2.04 \text{ nm}$  at the inlet and throat sections, respectively. The channel is periodic in both the streamwise and spanwise direction. The height between top and bottom walls  $h(x)$  varies in the streamwise direction according to a sinusoidal function,

$$h(x) = 2a \left[ \cos \left( \frac{2\pi x}{l} \right) - 1 \right] + h_{\text{inlet}}, \quad (5)$$

where  $4a = 1.36 \text{ nm}$  is the change of height from inlet to throat, and  $h_{\text{inlet}}$  is the height of the channel at the inlet.

The full MD domain is divided into 200 bins in the  $x$ -direction of bin-width  $\delta x = 0.34 \text{ nm}$ , and the instantaneous mass flow rate and density are measured in each bin. In the enhanced CFD domain, we define a plane across the channel at equivalent positions, and sum the mass fluxes from each cell the plane crosses, at each time-step, to get the instantaneous data. Dependency studies on the mesh resolution and on the time step showed that 50,000 cells and a time step of  $21.6 \text{ fs}$  were more than sufficient to obtain converged CFD solutions.

All the flows start from rest, then a time-varying gravity force  $F_g(t)$  is applied. We consider four different forces applied to the fluid:

1. **Startup flow:** a steady gravity force of  $F_g = 0.487 \text{ pN}$ .
2. **Short oscillations:** an unsteady, oscillating gravity force with amplitude  $0.487 \text{ pN}$  and period of  $T = 0.22 \text{ ns}$ , i.e.

$$F_g(t) = 0.487 \times 10^{-12} \sin \left( \frac{2\pi t}{0.22 \times 10^{-9}} \right), \quad (6)$$

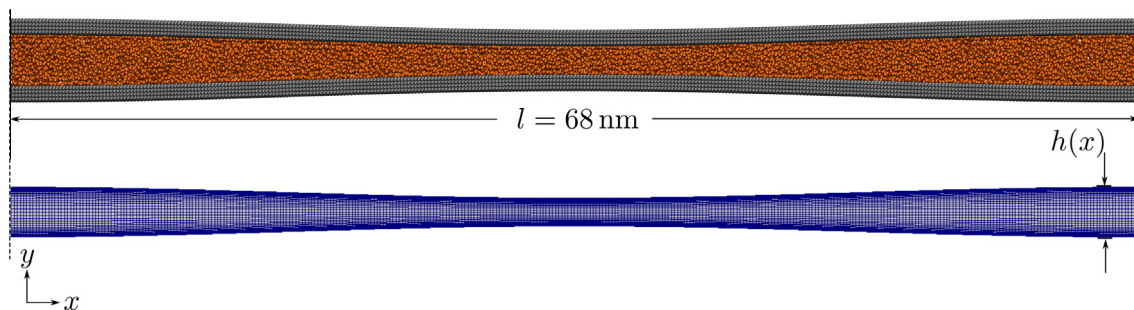
where  $t$  is the simulation time;

3. **Long oscillations:** an unsteady oscillating gravity force of the same amplitude, but with a larger period  $T = 10.8 \text{ ns}$ , i.e.

$$F_g(t) = 0.487 \times 10^{-12} \sin \left( \frac{2\pi t}{10.8 \times 10^{-9}} \right), \quad (7)$$

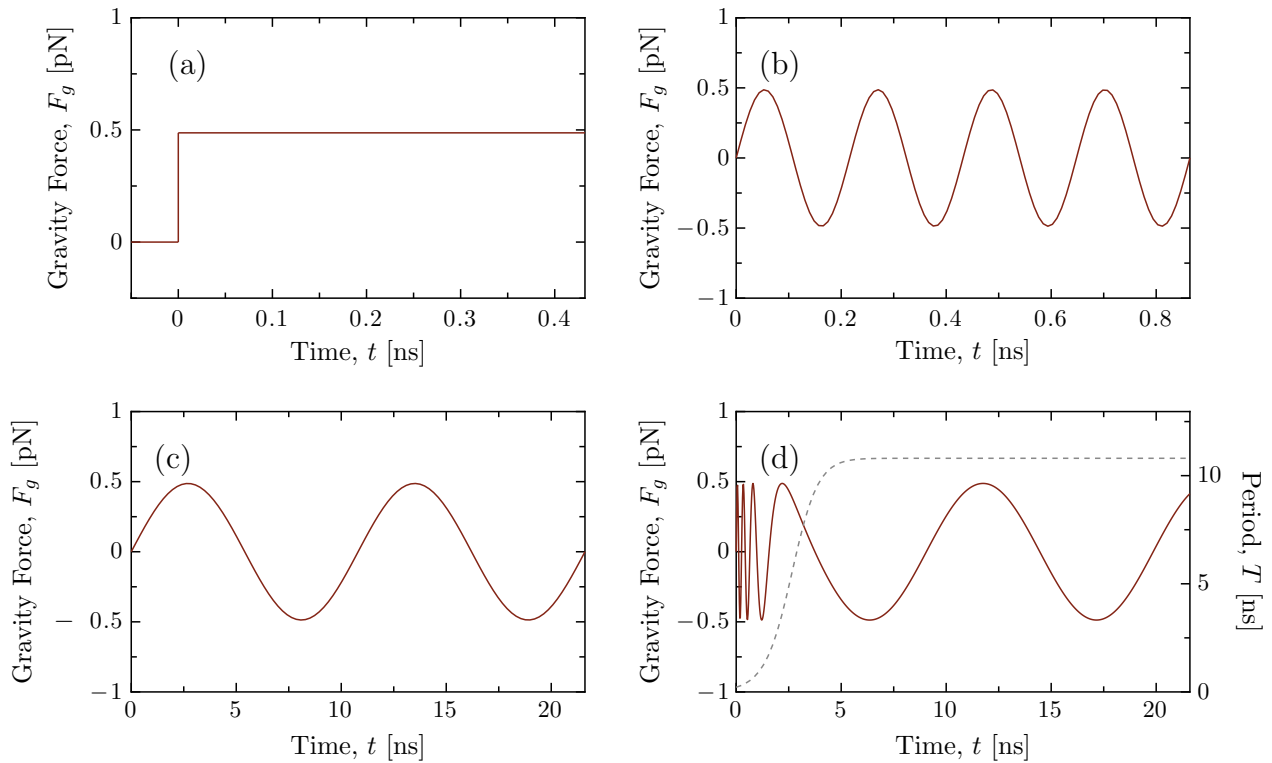
where  $t$  is the simulation time;

4. **Varying oscillations:** an unsteady oscillating gravity force with the same amplitude but with increasing period of  $0.2 \rightarrow 10.8 \text{ ns}$  as shown in Fig. 6(d), where the dashed line indicates how the period of the oscillation changes.



**Fig. 5.** The converging-diverging channel used in the unsteady flow cases. Top is the MD domain, and bottom is the CFD domain.





**Fig. 6.** The applied gravity forces varying with time for the four different cases: (a) step force, (b) oscillating gravity force with period  $T = 0.22$  ns, (c) oscillating gravity force with period  $T = 10.8$  ns, and (d) oscillating gravity force with increasing period  $T = 0.22$  ns  $\rightarrow$  10.8 ns, where the dashed line shows how the period of oscillation changes.

Graphical representations of how the forces vary are shown in Fig. 6.

### 3.2. Results for the four cases

To test the reliability of our CFD predictions that have MD pre-simulation input, we compare results with full-domain molecular dynamics calculations (referred to as ‘full MD’). To test whether our enhanced CFD model is an improvement over conventional CFD, we also compare results with predictions from compressible CFD with no-slip at the wall and without incorporating a CFD surface offset (referred to as ‘no-slip CFD’). We also compare with incompressible CFD with slip incorporated but no surface displacement (referred to below as ‘incomp. slip CFD’).

In Fig. 7 we plot the mass flow rate variation with time in a single bin near the inlet of the channel for each case. We see that in all cases the enhanced CFD model is able to accurately predict how the mass flow rate changes in time. Fig. 7(a), in which a constant force is applied throughout the channel, shows that the CFD reaches steady state at the same time as the MD simulation, and that a similar final mass flow rate is reached. There are, however, substantial differences between the enhanced CFD, the no-slip CFD, and the incomp. slip CFD results. The oscillations that are observed at the early times in Fig. 7(a) in the enhanced CFD results and also the MD data are due to an acoustic response of the nano channel to impulse forcing. A first estimate of the natural acoustic period is obtained by  $T = l/c = 0.07$  ns (where  $c$  is the speed of sound). This corresponds reasonably closely with the observed kinks in the mass flow rate.

Fig. 7(b) shows the results when an oscillating force with period 0.22 ns is applied. The mass flow rate in the enhanced CFD oscillates with the right frequency, the correct amplitude, and is also in phase with the full MD results. The no-slip CFD, on the other hand, appears to have the correct frequency but the amplitude is

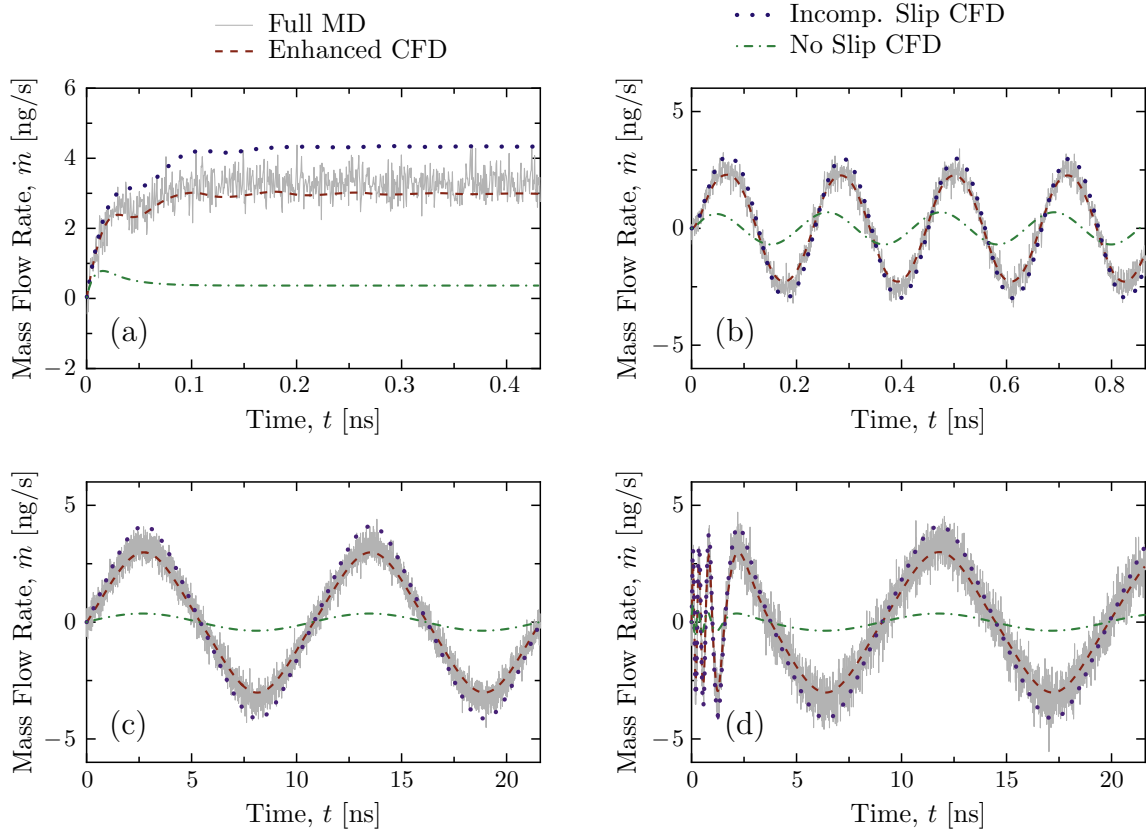
incorrect and it is oscillating out of phase, while the incomp. slip CFD is in phase but overpredicts the amplitude.

In Fig. 7(c) we have an oscillating force with period 10.8 ns. The mass flux in the enhanced CFD oscillates with the right frequency, correct amplitude, and is in phase, whereas the no-slip CFD appears to have the correct frequency, and is oscillating in phase, but the amplitude is incorrect. In Fig. 7(d) the period of the oscillating force increases from 0.22 ns to 10.8 ns; even in this more elaborate case, the enhanced CFD prediction is accurate.

Table 1 provides an indication of the computational cost for the full-domain MD simulations. The longest simulations presented in this paper ran in parallel (on 24 CPUs) for 48 days. The enhanced CFD itself has negligible computational cost by comparison, although the MD pre-simulations require the computational resources indicated in the last row of Table 1. However, these pre-simulations need only be performed once for a particular fluid/solid combination, and then can be used for any number of flow geometries thereafter.

## 4. Design optimisation

We now demonstrate how the enhanced CFD model can be used in design optimisation problems at the nanoscale. The example we choose is the optimal design of a bifurcating nano-channel network (see Fig. 8); such a design exploration would not be feasible using full MD simulations. The problem is to find the optimal widths of the channels in a bifurcating channel (i.e. those that give greatest mass flow rate), for a constant pressure difference  $\Delta p$  between the inlet and the outlets, and a constant volume  $V$ . At the macro scale the solution to this problem is given by Murray’s Law [29,30], which was first derived using the Hagen–Poiseuille Law to minimise the power required to sustain the flow of blood through vessels. It has also since been shown to describe the water transport through biological vessels in plants [31], and at the micro

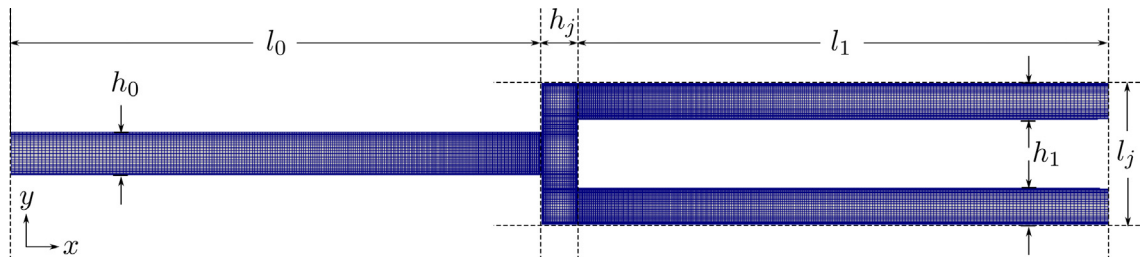


**Fig. 7.** The mass flow rate near the inlet of the channel varying with time, for each case. The solid lines are the full MD results, the dashed lines are the enhanced CFD results, the dotted lines are the incomp. slip CFD results and the dot dashed lines are the no-slip CFD results. (a) step force, (b) oscillating gravity force with period  $T = 0.22$  ns, (c) oscillating gravity force with period  $T = 10.8$  ns, and (d) oscillating gravity force with increasing period  $T = 0.22 \rightarrow 10.8$  ns. Note the statistical noise in the full MD results.

**Table 1**

Computational costs: the first four rows are for the full MD simulations, while the last row is the MD pre-simulation that is used to collect the data for the enhanced CFD.

	CPU	Liquid molecules	Wall molecules	Time per MD time-step (s)	Total computational time
Startup flow	24	69,264	19,677	0.68	16 h
Short oscillations	24	69,264	19,677	0.68	30 h
Long oscillations	24	69,264	19,677	0.68	48 days
Varying oscillations	24	69,264	19,677	0.68	48 days
MD pre-simulations	24	5073–6668	4160	0.14	4 days per liquid/solid combination



**Fig. 8.** The bifurcating channel domain used for the design optimisation. The width of the parent channel is  $h_0$ ; the width of the two daughter channels are  $h_1$ .

scale it has been used to optimally design MEMS devices with rectangular or trapezoidal cross sections [32].

For a 2D two-level network, like the geometry in Fig. 8, Murray's Law is

$$h_0^2 = \sum_{j=1}^N h_j^2, \quad (8)$$

where  $h_0$  is the width of the inlet parent channel, and  $h_1$  to  $h_N$  are the widths of the outlet daughter channels. For a symmetric bifurcating channel with  $N = 2$  and  $h_1 = h_2$ , the optimum ratio of channel widths is then given by

$$\frac{h_0^2}{2h_1^2} = 1. \quad (9)$$

The optimisation we perform, with the given constraints of constant volume and fixed pressure difference, is a linear search on channel width (equal increments) to find the maximum mass flow rate. We choose a volume of  $1100 \text{ nm}^3$ , a pressure difference of 10 MPa, channel lengths  $l_0 = l_1 = 75 \text{ nm}$ , a junction length  $l_j = 20 \text{ nm}$  and junction width,  $h_j = 4 \text{ nm}$ . The volume  $V$  can be calculated as:

$$V = h_0 l_0 + h_j l_j + 2h_1 l_1. \quad (10)$$

If this geometry was optimised using MD, each simulation would take approximately 30 days, whereas each enhanced CFD simulation takes approximately 500 s to perform. Fig. 9 shows the results from this optimisation with our enhanced CFD model used on a micro-scale channel and on a nano-scale channel. We see that for a micro scale channel, the optimum width occurs when  $h_0^2/2h_1^2 = 1$ : this is the expected result according to Murray's Law. At the nano-scale, however, we observe a significant deviation from the standard Murray's Law, which is now discussed.

A deviation from the standard Murray's Law has been noted for rarefied gases [33] but has not so far been demonstrated for a liquid. To uncover the origin of this deviation we derive Murray's Law using Poiseuille's equations with Navier slip at the walls, i.e.  $u(h) = u(-h) = \xi \frac{du}{dy}$  where  $\xi$  is the slip length, and the velocity is at a maximum at  $y = 0$  i.e.  $\frac{du}{dy}|_{y=0} = 0$ . The mass flow rate is then

$$\dot{m} = \frac{2\rho}{3\mu} \frac{\Delta p}{l} h^3 \left(1 + \frac{3\xi}{h}\right), \quad (11)$$

where  $\dot{m}$  is the mass flow rate,  $\rho$  is the density,  $\mu$  is the dynamic viscosity and  $\Delta p$  is the pressure difference between the inlet of the parent channel and the outlet of the daughter channel. Murray's Law is found by minimising the power  $P$  required to maintain flow, which for flow through a channel is

$$P = \dot{m}\Delta p + 2bhl, \quad (12)$$

where  $b$  is a constant of proportionality. By eliminating  $\Delta p$  with Eq. (11) in this equation and differentiating, we find that when the power is minimised the mass flow rate is

$$\dot{m} = kh^2 \frac{(1 + 3\xi/h)}{(1 + 2\xi/h)^{1/2}}, \quad (13)$$

where  $k = 2/3\sqrt{\rho b/\mu}$ . For a symmetric bifurcating channel, the mass flow rate through the parent channel must equal the total mass flow rate through the daughter channels, i.e.  $\dot{m}_0 = 2\dot{m}_1$ , therefore, the optimal ratio of channel widths becomes

$$\frac{h_0^2}{2h_1^2} = \frac{(1 + 3\xi/h_1)}{(1 + 3\xi/h_0)} \frac{(1 + 2\xi/h_0)^{1/2}}{(1 + 2\xi/h_1)^{1/2}}. \quad (14)$$

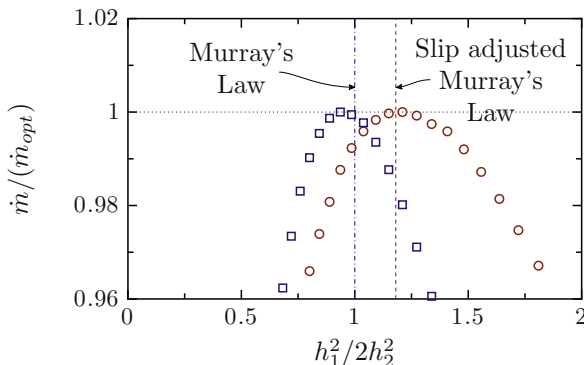


Fig. 9. The normalised mass flow rate for different ratios of mother/daughter channels widths for micro-scale channels (squares) where the dash-dotted line is the expected optimum, and for nano-scale channels (circles) where the dashed line is the expected optimum.

It is clear that when  $h_0, h_1 \gg \xi$  this becomes Eq. (9), as expected. It can also be noted that when the flow becomes plug-like, i.e.  $h_0, h_1 \ll \xi$ , this ratio becomes  $h_0^2/2h_1^2 = 2^{1/3}$ .

We can now use Eqs. (10) and (14) to calculate the expected value of  $h_0^2/2h_1^2$ . When comparing this to the optimum found by the enhanced CFD we get excellent agreement, as highlighted in Fig. 9. This shows that the slip at the walls is the important factor in the deviation from the expected optimum. A CFD model that includes an accurate model of the wall–fluid interaction is, therefore, potentially very important in the design of nano-scale devices.

## 5. Summary

We have shown that a CFD model enhanced with data from MD pre-simulations is capable of making accurate predictions of unsteady liquid flow along a converging–diverging channel that has a width close to the expected continuum–fluid limit. This enhanced CFD approach is far more accurate than conventional CFD calculations, and significantly more computationally efficient than full MD simulations.

We have also demonstrated the enhanced CFD approach applied to a design optimisation problem: that of a bifurcating nanofluidic network. The widths of channels in the network should be optimised to maximise the mass flow rate through the network, for a fixed pressure drop and network volume. We have shown that slip at the nano-scale can have a very significant effect on the optimum channel dimensions, and we have derived an analytical equation which corrects the well-known Murray's Law. This is one of many possible cases where nano-scale flow effects modify the optimal design of nanofluidic systems when compared with their macroscopic counterparts.

## Acknowledgments

This work is financially supported in the UK by EPSRC Programme Grant EP/I011927/1 and EPSRC Grants EP/K038664/1 and EP/K038621/1. Our calculations were performed on the high performance computer ARCHIE at the University of Strathclyde, funded by EPSRC Grants EP/K000586/1 and EP/K000195/1.

## References

- [1] Alexiadis A, Kassinos S. Molecular simulation of water in carbon nanotubes. *Chem Rev* 2008;108(12):5014–34.
- [2] Mantzalis D, Asproulis N, Drikakis D. Filtering carbon dioxide through carbon nanotubes. *Chem Phys Lett* 2011;506(1):81–5.
- [3] Saidur R, Leong KY, Mohammad HA. A review on applications and challenges of nanofluids. *Renew Sustain Energy Rev* 2011;15:1646–68.
- [4] Wen D, Lin G, Vafaei S, Zhang K. Review of nanofluids for heat transfer applications. *Particuology* 2009;7(2):141–50.
- [5] Levitt M, Warshel A. Computer simulation of protein folding. *Nature* 1975;253(5494):694–8.
- [6] Parrinello M, Rahman A. Crystal structure and pair potentials: a molecular-dynamics study. *Phys Rev Lett* 1980;45(14):1196.
- [7] Sheehan ME, Sharratt PN. Molecular dynamics methodology for the study of the solvent effects on a concentrated diels-alder reaction and the separation of the post-reaction mixture. *Comput Chem Eng* 1998;22:S27–33.
- [8] Reese JM, Gallis MA, Lockerby DA. New directions in fluid dynamics: non-equilibrium aerodynamic and microsystem flows. *Philos Trans Royal Soc Lond Series A: Math, Phys Eng Sci* 2003;361(1813):2967–88.
- [9] Bocquet L, Charlaix E. Nanofluidics, from bulk to interfaces. *Chem Soc Rev* 2010;39(3):1073–95.
- [10] Huang C, Choi PY, Nandakumar K, Kostiuk LW. Comparative study between continuum and atomistic approaches of liquid flow through a finite length cylindrical nanopore. *J Chem Phys* 2007;126(22):224702.
- [11] Sofos F, Karakasidis T, Liakopoulos A. Transport properties of liquid argon in krypton nanochannels: anisotropy and non-homogeneity introduced by the solid walls. *Int J Heat Mass Transf* 2009;52(3–4):735–43.
- [12] Travis KP, Todd BD, Evans DJ. Departure from Navier–Stokes hydrodynamics in confined liquids. *Phys Rev E* 1997;55(4):4288.
- [13] Todd BD, Hansen JS, Daivis PJ. Nonlocal shear stress for homogeneous fluids. *Phys Rev Lett* 2008;100(19):195901.



- [14] Cadusch PJ, Todd BD, Zhang J, Daivis PJ. A non-local hydrodynamic model for the shear viscosity of confined fluids: analysis of a homogeneous kernel. *J Phys A: Math Theor* 2008;41(3):035501.
- [15] Todd BD. Cats, maps and nanoflows: some recent developments in nonequilibrium nanofluidics. *Mol Simul* 2005;31(6-7):411–28.
- [16] Holland DM, Lockerby DA, Borg MK, Nicholls WD, Reese JM. Molecular dynamics pre-simulations for nanoscale computational fluid dynamics. *Microfluid Nanofluid*. <http://dx.doi.org/10.1007/s10404-014-1443-6>.
- [17] Popadić A, Walther JH, Koumoutsakos P, Praprotnik M. Continuum simulations of water flow in carbon nanotube membranes. *New J Phys* 2014;16(8):082001.
- [18] Mohamed KM, Mohamad AA. A review of the development of hybrid atomistic–continuum methods for dense fluids. *Microfluid Nanofluid* 2010;8(3):283–302.
- [19] Borg MK, Lockerby DA, Reese JM. The FADE mass-stat: a technique for inserting or deleting particles in molecular dynamics simulations. *J Chem Phys* 2014;140(7):074110.
- [20] Borg MK, Macpherson GB, Reese JM. Controllers for imposing continuum-to-molecular boundary conditions in arbitrary fluid flow geometries. *Mol Simul* 2010;36(10):745–57.
- [21] Macpherson GB, Borg MK, Reese JM. Generation of initial molecular dynamics configurations in arbitrary geometries and in parallel. *Mol Simul* 2007;33(15):1199–212.
- [22] Macpherson GB, Reese JM. Molecular dynamics in arbitrary geometries: parallel evaluation of pair forces. *Mol Simul* 2008;34(1):97–115.
- [23] OpenFOAM. The open source CFD toolbox; 2015. <<http://www.openfoam.org>>.
- [24] Thompson PA, Troian SM. A general boundary condition for liquid flow at solid surfaces. *Nature* 1997;389(6649):360–2.
- [25] Irving JH, Kirkwood JG. The statistical mechanical theory of transport processes. iv. The equations of hydrodynamics. *J Chem Phys* 1950;18:817–29.
- [26] Linstrom PJ, Mallard WG. NIST chemistry webbook. NIST standard reference database no. 69. <<http://webbook.nist.gov>>.
- [27] Gad-el Hak M. MEMS: introduction and fundamentals. CRC Press; 2010.
- [28] Patronis A, Lockerby DA, Borg MK, Reese JM. Hybrid continuum–molecular modelling of multiscale internal gas flows. *J Comput Phys* 2013;255:558–71.
- [29] Murray CD. The physiological principle of minimum work: I. The vascular system and the cost of blood volume. *Proc Natl Acad Sci USA* 1926;12(3):207–14.
- [30] Murray CD. The physiological principle of minimum work: II. Oxygen exchange in capillaries. *Proc Natl Acad Sci USA* 1926;12(5):299–304.
- [31] McCulloh KA, Sperry JS, Adler FR. Water transport in plants obeys Murray's law. *Nature* 2003;421(6926):939–42.
- [32] Emerson DR, Cieřlicki K, Gu X, Barber RW. Biomimetic design of microfluidic manifolds based on a generalised Murray's law. *Lab Chip* 2006;6(3):447–54.
- [33] Gosselin L, da Silva AK. Constructal microchannel networks of rarefied gas with minimal flow resistance. *J Appl Phys* 2007;101(11):114902.